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1 *Article*

# 2 **Clinical characteristics and management of** 3 **neurocysticercosis patients: a retrospective assessment** 4 **of case reports from Europe**

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52 **Abstract:**

53 **Objectives:** Neurocysticercosis (NCC) is a parasitic disease caused by the larval stage of the tapeworm *Taenia*  
54 *solium*. NCC mainly occurs in Africa, Latin America and South-East Asia and can cause a variety of clinical  
55 signs/symptoms. Although it is a rare disease in Europe, it should nonetheless be considered as a differential  
56 diagnosis. The aim of this study was to describe clinical characteristics and management of patients with NCC  
57 diagnosed and treated in Europe.

58 **Methods:** We conducted a systematic search of published and unpublished data on patients diagnosed  
59 with NCC in Europe (2000–2019) and extracted demographic, clinical and radiological information on each case, if  
60 available.

61 **Results:** Out of 293 identified NCC cases, 59% of patients presented initially with epileptic seizures (21%  
62 focal onset); 52% presented with headache and 54% had other neurological signs/symptoms. The majority of  
63 patients had a travel or migration history (76%), mostly from/to Latin America (38%), Africa (32%) or Asia (30%).  
64 Treatment varied largely depending on cyst location and number. The outcome was favorable in 90% of the cases.

65 **Conclusions:** Management of NCC in Europe varied considerably but often had a good outcome. Travel and  
66 migration to and from areas endemic for *T. solium* will likely result in continued low prevalence of NCC in Europe.  
67 Therefore, training and guidance of clinicians is recommended for optimal patient management.

68

69 **Keywords:** Neurocysticercosis; *Taenia solium*; Europe; neglected tropical diseases; NCC management; Global  
70 Health; Clinical epidemiology; One Health

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74 **INTRODUCTION**

75 Neurocysticercosis (NCC) is caused by the tapeworm *Taenia solium*, a zoonotic parasite which has a pig-  
76 human-environment life cycle. Humans get NCC by ingesting parasite eggs, which are shed through  
77 feces of *T. solium* tapeworm carriers. Through environmental contamination and lack of hygiene these  
78 eggs can accidentally be ingested by humans. Once in the intestines, the larvae, which are released from  
79 eggs, cross the intestinal mucosa into the circulatory system through which they are transported to  
80 multiple organs, where they encyst (cysticerci). When these cysticerci affect the central nervous system  
81 (CNS), the disease is called NCC. Cysts can persist in the CNS for several years without causing any  
82 neurological signs/symptoms. Signs and symptoms are pleomorphic [1–4], often resulting from  
83 degeneration of the cysts and associated inflammatory host immune reaction. The most common  
84 neurological signs/symptoms are epileptic seizures, headache episodes, focal neurological deficits and  
85 signs of raised intracranial pressure [1].

86 Neurological signs and symptoms depend on number, size, location (e.g. intraparenchymal or  
87 extraparenchymal and location within the brain parenchyma), and cyst stage. When located in the  
88 parenchyma, degeneration of cysts (colloidal and granular nodular stage) is associated with  
89 inflammation leading to perilesional oedema which can cause neurological signs/symptoms [2,5].

90 Treatment options for intraparenchymal lesions include anthelmintic therapy with albendazole and/or  
91 praziquantel in combination with corticosteroids for vesicular cysts, or corticosteroids alone for  
92 degenerating cysts; both accompanied by anti-epileptic drugs, if necessary. For extraparenchymal  
93 lesions treatment options include ventriculoperitoneal shunting, or surgical removal of cysts [4,6–13].

94 According to the distribution map for NCC published by WHO, Latin America, South and South-East  
95 Asia and sub-Saharan Africa are considered endemic [14]. In these areas, NCC accounts for around one  
96 third of all epilepsy cases [4,15]. In Europe, although data are scarce, the main countries affected are  
97 Spain, Portugal and Eastern European countries, however only rarely [16–21]. Until the 1990s, many  
98 autochthonous cases of *T. solium* infection were reported in Portugal and Spain.[16] To date, many  
99 immigrants who are likely to have been infected outside Europe are also diagnosed in these two  
100 countries. [16] In Eastern Europe, disease surveillance is only sporadic; a previous review reported a  
101 particularly large number of cases from Serbia with probable infection in Eastern Europe.

102 Disease presentation differs between world regions. While single enhancing intraparenchymal lesions  
103 are the predominant form of the disease in India, multiple lesions are common in Latin America which  
104 more commonly are also in the extraparenchymal space than in other world regions. For African  
105 populations, also multiple lesions which are mostly located in the parenchyma have been described  
106 [2,22–25]. For Europe, disease presentation has not yet been described.

107 Knowledge of diagnostic work-up and management of patients presenting with symptomatic NCC may  
108 be scarce among clinicians in Europe where NCC cases are rare and most clinicians have never seen a

109 NCC case [26]. As knowledge about differences of disease manifestation due to geographical  
110 characteristics or certain risk factors may help to correctly diagnose and treat patients, the aim of this  
111 study was to summarize and update information about clinical characteristics and management of NCC  
112 patients diagnosed in Europe.

## 113 **METHODS**

114 **Systematic literature search:** This project was part of CYSTINET (European Network on  
115 Taeniosis/Cysticercosis, COST Action TD 1302) [27]. We conducted a systematic literature search of  
116 NCC case reports and case series, along with cases from the grey literature. Also reviewed were  
117 unpublished data collected via collaborating clinicians and laboratories. Moreover, experts familiar  
118 with NCC patient management in the European setting were consulted. The protocol for the conduct of  
119 systematic literature review followed the PRISMA-P outline and was registered on PROSPERO  
120 (registration number: CRD42016050729) [28]. Ethical approval was obtained where required. Ethical  
121 approval for the retrospective analysis of anonymized patient data was granted by the ethics committee  
122 of the Klinikum rechts der Isar at the Technical University of Munich, Germany (208/16S).

123 **Search methods:** PubMed, EMBASE, Web of Science, Global Health (CABI), Global Index Medicus  
124 coupled with Aoister and Open Grey were searched for articles published between January 2000 to May  
125 2019. Supplement Table S1 contains the precise search terms and dates. Moreover, each CYSTINET  
126 researcher searched for grey literature in their native countries (25 countries; the list of countries can be  
127 found under this link: <http://www.cystinet.org/the-action/participating-countries/>); references of  
128 included literature were evaluated for relevance and included if they met the inclusion criteria. No  
129 language restriction was applied; only studies on humans were included. (Systematic) reviews were  
130 screened for additional references [16,17,19].

131 **Study selection criteria:** Inclusion and exclusion criteria were pre-defined in the study protocol  
132 (Supplement Table S2). Only data on NCC cases presenting in Europe (see list of included countries in  
133 Supplement Table S2) were included. Case reports and case series were considered for inclusion. NCC  
134 was defined as the presence of *T. solium* cysts/calcifications in the CNS confirmed on neuroimaging.  
135 Studies were excluded if 1) reporting on another *Taenia* species (e.g. *T. crassiceps*, *T. hydatigena*, *T.*  
136 *asiatica*), 2) reporting on cysticercosis only outside the CNS (e.g. muscles, eyes etc.), 3) reporting on  
137 patients treated outside Europe, 4) reporting year before 2000 (even if published after 2000), 5) reporting  
138 on the same patients (when reporting on the same patient both articles were taken into account for  
139 additional information, but the patient was counted as one) and 6) reporting on animals.

140 **Study selection process:** The Covidence online tool (<https://www.covidence.org/>) was employed to  
141 assess the published literature obtained from PubMed, EMBASE, Web of Science, Global Health (CABI),  
142 and Global Index Medicus [29]. Literature was screened independently by four reviewers (AA, JB, PS,

143 CU). Each selection required two votes from the reviewers; in the event of disagreement, a third  
144 reviewer was consulted. First, the titles and abstracts were screened and a decision was made whether  
145 to include or exclude the publication.

146 Next, publications were sorted by country name (Supplement Table S2), and if found suitable as per the  
147 inclusion criteria, retained. Following this, the complete texts of the papers were reviewed and the  
148 rationale for any exclusion specified. In order to confirm the validity and suitability of the  
149 inclusion/exclusion criteria, the procedure of study selection was test-run by all researchers. Searching  
150 Aoister and Open Grey in collaboration with CYSTINET members yielded grey literature, including  
151 doctoral theses, papers in languages other than English and conference abstracts [17,19], based on the  
152 same selection process as described above.

153 Collection of unpublished data: Attendees at the 3-4 November 2015 CYSTINET international  
154 conference in Belgrade, Serbia, were surveyed by means of a questionnaire in order to obtain further  
155 grey literature and unpublished data. Details of patient data - with due regard to in-country ethical  
156 stipulations – as well as information sources and local experts' contacts were also requested from the  
157 CYSTINET members. Three of the authors (EH, NFW, PLC) collated the primary source data for a series  
158 of 26 cases managed at Hospital of Tropical Diseases in London, United Kingdom, that were  
159 subsequently published after the present study was conceived and initiated [30]. Those cases are  
160 described here as unpublished, which accurately reflects their status at the time the data for the present  
161 study were collated. Ethical approval was obtained where required (Serbia, Portugal, Romania). At all  
162 subsequent CYSTINET meetings and conferences, reminders were issued and contributions were also  
163 solicited via email. Medical plausibility verifications were conducted (by AA, DS, MK and ASW) on the  
164 patient data, which were all anonymized.

165 **Data extraction:** All variables for data extraction have been outlined in the research protocol; these were  
166 consequently utilized in the data extraction process. Data were extracted by five independent  
167 researchers (DS, RM, AA, AF, MK) and in case of uncertainty another expert of the group (ASW) was  
168 consulted. Plausibility verifications were carried out on the data extracts saved in Excel sheets, by a  
169 different reviewer from the one who had extracted the data.

170 **Definition of variables:** Autochthonous cases were defined as not having migrated from or never  
171 having travelled to an area outside of Europe that is endemic for *T. solium*. Only if travel/migration  
172 history was specifically denied by the patient, was the case considered to be autochthonous; otherwise  
173 the information was considered to be not available. Epileptic seizure types were classified as focal onset  
174 or generalised onset seizures according to the latest International League against Epilepsy (ILAE)  
175 definition [31]. For the evaluation of diagnostic variables computed tomography (CT), magnetic  
176 resonance imaging (MRI), soft tissue x-ray, and electroencephalography (EEG) were recorded.  
177 Furthermore, variables on location (cerebral: intraparenchymal, intraventricular, subarachnoid; spinal:

178 intra-medullary/extra-medullary, and extra-neural) and stages of the cyst(s) (active: vesicular, colloidal,  
179 granular nodular; inactive: calcified) were extracted. In addition, other diagnostic findings such as  
180 perilesional edema and hydrocephalus were noted. All information mentioned in the text or visible on  
181 pictures was included. Favourable outcomes were defined as “Cured” or “Improved”. If patients were  
182 free from symptoms and no active cysts were visible on follow-up imaging, the patient was considered  
183 as cured. Also, if it was specifically mentioned that the patient was cured. The patient was considered  
184 to have improved if at least one active cyst shrank in size or if symptoms after treatment were less  
185 intense or less frequent as before.

186 **Statistical analyses:** Categorical variables were compared with Chi-square tests and Chi-square tests  
187 for trends where applicable. Continuous variables were compared using the Wilcoxon test when non-  
188 normally distributed. Statistical analyses were performed using R version 3.6.2 [32].

189

## 190 **RESULTS**

### 191 *Search Results*

192 Searching PubMed, EMBASE, Web of Science, Global Health (CABI), and Global Index Medicus  
193 identified a total of 13,264 publications. Through Aoister, Open Grey, CYSTINET presentations and  
194 through personal communication 52 additional publications were found. After de-duplication, 10,088  
195 remained. After title, abstract and full text screening, 145 publications on individual cases or smaller  
196 case series (containing data of overall 211 patients with NCC) were included. The search process is  
197 presented in a flowchart in Supplement Figure S1. Through expert consultations we retrieved a further  
198 82 unpublished cases of NCC. Most of the unpublished cases were from the United Kingdom (n=34,  
199 41%) and Romania (n=31, 38%), but we also received case descriptions from Austria, France, Germany  
200 and Italy. More than 50% of the published cases were diagnosed and treated in three western European  
201 countries, namely Spain (48/211), France (38/211) and Portugal (28/211). A further 16 countries also  
202 reported cases, of which four countries reported more than 10 cases (United Kingdom, Germany, Italy  
203 and Slovenia; Figure 1, Supplement Table S3).

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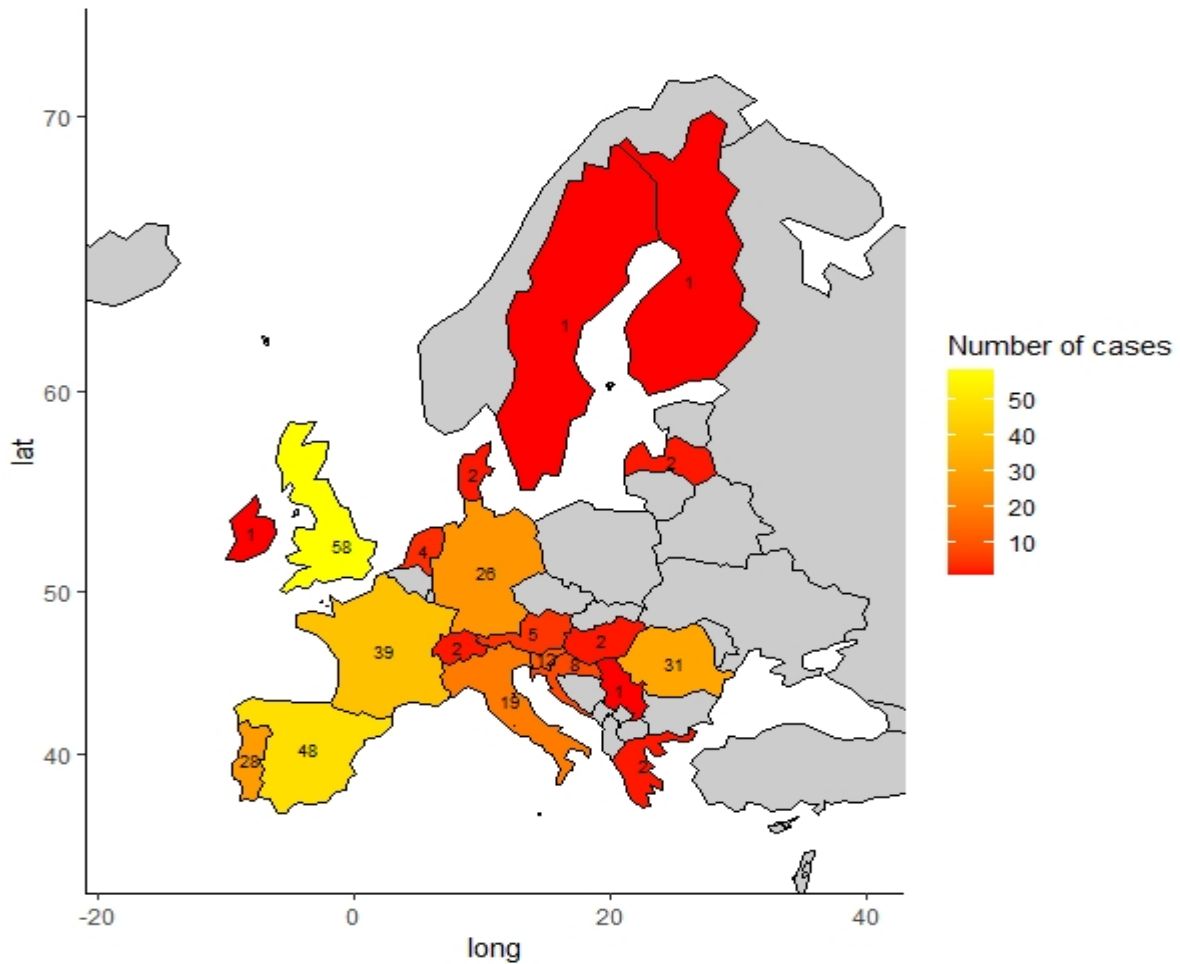
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219 **FIGURE 1.** Map of Europe showing the number of cases included in this analysis

220

221 *Patient demographic and clinical characteristics*

222 *Demographics and migration details*

223 Among the published cases, 53% of those with recorded sex, were female. Median age at diagnosis was  
224 32 years (interquartile range 21 to 47); the youngest patient was two years old and the oldest patient  
225 was 82 years old. Nearly every fifth (n=38, 19%) patient was a child or an adolescent (aged <18 years;  
226 Supplement Figure S2). Most patients (85%) either originated from or travelled to areas highly endemic  
227 for *T. solium*. Only 15% of all patients denied having travelled outside Europe; these patients were  
228 considered autochthonous cases (Table 1). More than half of those cases either occurred in or had a  
229 travel/migration history to/from eastern European countries, e.g. Hungary or Romania. Among the



230 “imported” cases, approximately half of all patients migrated from or had travelled to Latin America,  
231 mostly from Ecuador, Colombia, Bolivia or Brazil. Most of these patients were treated in Spain (61%).  
232 Most cases from Asia had a travel history from India (22/33); most cases from Africa were diagnosed in  
233 Portugal, the majority originated from Cape Verde (17/44) and Guinea-Bissau (5/44). A substantial  
234 proportion of unpublished cases were reported from Romania where most patients had never left  
235 Europe or even their country. Hence, almost 50% (32/70) of the unpublished cases were autochthonous  
236 cases. Contrary to the published cases, among unpublished cases with travel/migration history, most  
237 migrated/travelled from Asia (58%) or Africa (42%).

238

#### 239 *Neuroimaging and electroencephalography*

240 All patients reported in this paper had neuroimaging performed, but it was not always specified  
241 whether this was a CT and/or an MRI. In the majority of cases, diagnostic imaging was performed with  
242 a combination of CT and MRI, usually including contrast medium; in only 16% of the cases, diagnosis  
243 was based solely on CT scanning. In approximately one quarter of patients soft tissue imaging was also  
244 performed. Indications for this were most often either extensive findings on neuroimaging or palpable  
245 rice corn like cysts/calcifications on clinical examination. Among the published cases, 24 (14%) had an  
246 EEG documented (15 with result); of those, 4 had a normal EEG, 3 showed epileptic activity and 8  
247 showed abnormal unspecified patterns. Those were mostly patients with epileptic seizures.

248

#### 249 *Staging of cysts*

250 Around 90% of all NCC patients (published 89% and unpublished 93%; Table 1) had viable cysts in their  
251 brain (most commonly in the parenchyma) or spine. Viable cysts were defined as cysts in vesicular or  
252 degenerative stage. The remaining patients presented with calcifications only. There is some indication  
253 that children more often had intraparenchymal NCC (26/29, 90%), whereas adults scored higher on  
254 intraventricular (39/199, 20%) and subarachnoid NCC (25/197, 13%; Supplement Table S4). Ninety-one  
255 (41%) patients had only a single lesion (regardless of whether viable or calcified; Table 1). Of the patients  
256 with viable cysts, published and unpublished cases taken together, 108 patients (68%) had at least one  
257 cyst with ring enhancement and 85 (59%) had a scolex visible on imaging. Perilesional edema was  
258 present in 124 patients (57%; Table 1). There was one case report of a patient who only had one  
259 calcification – no viable cysts – but perilesional edema. Thirty-eight patients had hydrocephalus, most  
260 of which were patients with extraparenchymal or spinal cysts. Five patients with hydrocephalus were  
261 described as having only intraparenchymal cysts. Among the published cases, the majority of patients  
262 had at least one cyst in the degenerative (colloidal or granular nodular) stage (57%). Twenty-one of the

263 144 of the patients with available detailed information on the stage of the cysts had calcifications only  
264 (Table 1). Five of 27 patients with subarachnoid cysts also had spinal cysts.

265

#### 266 *Radiological differences by origin of infection*

267 Patients who migrated from Latin America had the highest proportion of extraparenchymal NCC, either  
268 only extraparenchymal lesions or in combination with intraparenchymal lesions (47%), and they mostly  
269 had multiple lesions (62%). Those migrating from Asia and Africa had the smallest proportion of  
270 extraparenchymal (26%) and multiple lesions (42%), respectively. Presentation of European  
271 autochthonous cases were in-between the other world regions (Supplement Tables S5/6). Patients with  
272 infection Latin America only seldomly had extraneural lesions (3/50; 6%). This proportion was highest  
273 for patients with autochthonous European infections (8/21; 38%).

274

#### 275 *Neurological signs/symptoms*

276 The most common presentation in both the published (55%) and the unpublished (72%) cases, were  
277 epileptic seizures. Children more commonly presented with epileptic seizures than adults (Supplement  
278 Table S4). Most of the seizures were of generalised onset. Concomitantly, many patients reported  
279 headache episodes (52%). Twenty-three patients presented only with headache (12%) without any other  
280 neurological signs/symptoms. More than half of all patients (54%) presented with other neurological  
281 signs/symptoms either alone or in combination with headache and/or epileptic seizures (Table 1). Those  
282 signs/symptoms were most commonly unsteady gait (27%), cognitive impairment (21%), impaired  
283 consciousness (20%) or impaired vision (19%). Also reported were cases with cranial nerve lesions,  
284 speech difficulties, meningism and vertigo (Table 2).

285

#### 286 *Association of neurological signs/symptoms and neuroimaging results*

287 Neurological signs/symptoms varied depending on cyst location, number, stage and other additional  
288 findings (Table 3, Figure 2, Supplement Figure S3). Regarding cyst location, presentation with epileptic  
289 seizures was more common in patients with intraparenchymal cysts. Epileptic seizures were reported  
290 in 66% of patients with intraparenchymal cysts compared with 25% of patients with at least one  
291 intraventricular cyst and 36% of patients with at least one subarachnoid cyst. This difference was even  
292 more pronounced when excluding patients with cysts at various locations (72% versus 15%/14%,  
293  $p < 0.005$ ; Table 3 and Figure 2A). Patients with only intraventricular cysts were more likely to present  
294 with headache than patients with only intraparenchymal or subarachnoid cysts (86% versus 43%/57%;  
295  $p < 0.01$ ; Table 3). Furthermore, patients with subarachnoid cysts, were significantly older than patients  
296 with cysts at other locations (median: 47 years [IQR 32–56 years] versus 31 years [IQR 24–45]; Wilcoxon

297 test  $p < 0.005$ , Supplement Figure S4). The majority of patients who showed hydrocephalus also had  
298 headache (67% versus 46%,  $p = 0.04$ ) or other neurological signs/symptoms (85% versus 34%,  $p < 0.005$ ;  
299 Table 3 and Figure 2B).

300 Regarding cyst stage, patients with cysts in the vesicular stage more commonly presented with  
301 headache compared to degenerative and calcified cyst stage (65% versus 44%/38%;  $p = 0.06$ ). Patients  
302 with degenerative (71%) or calcified (65%) cyst stage more commonly presented with epileptic seizures  
303 ( $p < 0.005$ ; Table 3, Figure 2C). There was no significant difference in the number of patients with other  
304 neurological signs/symptoms between the cyst stage (viable/degenerative/calcified: 47%/48%/42%,  
305  $p = 0.84$ ).

306 With respect to cyst number, patients with single lesions more commonly presented with seizures than  
307 patients with multiple lesions ( $p < 0.005$ ; Table 3), but there was no difference for headache or other  
308 neurological signs/symptoms. Among the published cases, seven had spinal cysts only. Symptoms  
309 ranged from general disorientation and headache, which could indicate undetected cerebral  
310 involvement, to brachialgia, brachial paralysis, bladder dysfunction, L5 radiculopathy, steppage gait  
311 and cauda equina syndrome. Overall, 30 patients additionally had extra-neural lesions. Most common  
312 locations were ocular cysts ( $n = 8$ ), cysts in the thoracic/back muscles ( $n = 10$ ) and calcifications in the thigh  
313 muscles ( $n = 7$ ).

314

#### 315 *Laboratory tests*

316 As determined by the inclusion criteria, all patients had NCC confirmed on neuroimaging. In addition,  
317 one hundred and eighty-four patients (63%) had serological testing of which 131 (71%) were positive in  
318 any test (serum or CSF, antigen or antibody). Among published cases 73% had a positive test, of the  
319 unpublished cases only 68% tested positive (Table 1). For those with information on diagnostic tests  
320 available, eight patients were reported to have been tested for antigen (three serum and CSF, four only  
321 serum, one only CSF); seven of these patients were antigen positive, one had an indeterminate result.  
322 Seventy-one patients were tested for *T. solium* specific antibodies (37 serum and CSF, 34 only serum and  
323 0 only CSF) and 61 (86%) were positive in any test. Western blot was more commonly used than ELISA  
324 in both serum and CSF (Supplement Table S7). Patients with single lesions were less commonly positive  
325 in any serological test than patients with multiple lesions (46% versus 75%, Supplement Table S8). Also,  
326 patients with extraparenchymal lesions more commonly were serologically positive (Supplement Table  
327 S8). Thirty patients had stool examined of which none was positive for *T. solium* eggs.

328

#### 329 *Treatment and outcomes*

330 Anthelmintic therapy was used in the treatment of the majority of NCC cases: 191 patients (81%) were  
 331 treated with anthelmintics. Most patients received albendazole, either alone (76%) or in combination  
 332 with praziquantel (15%; Table 1). The duration of anthelmintic treatment ranged from a single dose to  
 333 three months (praziquantel) and from a single dose to nine months (albendazole). The most common  
 334 treatment duration was 10–15 days (Supplement Figure S5), longer for patients with extraparenchymal  
 335 lesions compared to those with intraparenchymal lesions (Supplement Table S9). Eleven patients with  
 336 only extraparenchymal lesions were treated with anthelmintic medication, e.g. after extirpation of  
 337 spinal cysts.

338 One-hundred-fifty-five patients (73%) received corticosteroid therapy, either dexamethasone or  
 339 prednisolone/prednisone. Of those, 147 (95%) patients were also treated with anthelmintics, and 8 (5%)  
 340 patients were treated with steroids alone. Dexamethasone was more frequently used than  
 341 prednisolone/prednisone (68% versus 41%; some used both; Table 1).

342 Surgical treatment was performed in 71 patients (40%), mainly patients with intraventricular cysts.  
 343 Surgical treatment often involved extirpation of the cysts and drainage of the hydrocephalus through  
 344 ventricular shunting. Extirpated cysts were usually analysed pathologically (Table 1).

345 The majority of patients presenting with epileptic seizures were put on antiepileptic drugs (AED) unless  
 346 they had been on AED already. Twenty-five patients (21%) did not receive AED despite presenting with  
 347 epileptic seizures. The most common AED was carbamazepine (400mg/d), followed by levetiracetam  
 348 (1000mg/d) and valproic acid (1000mg/d; Supplement Table S10).

349 When reported, the treatment outcome was favourable in 139/155 (90%) patients, although only 68 (44%)  
 350 patients were reported to have been cured from NCC. Ten percent of the patients did not have an  
 351 improvement of symptoms or lesions – some even deteriorated and five patients, of which four were  
 352 younger than 40 years old, died from the disease during or after treatment (Table 1). Two of the patients  
 353 who died had intraventricular cysts, one patient concomitantly had a glioblastome multiforme, one  
 354 patient was living with HIV and developed bronchopneumonia during therapy, and the fifth patient  
 355 died from epileptic seizures. Supplement Table S11 shows treatment outcome by various parameters.  
 356 The outcome did not differ significantly for any of the parameters ( $p>0.05$  for all). Children more  
 357 commonly were cured through therapy than adults (20/35 [57%] versus 48/120 [40%]).

358

359 **TABLE 1.** Characteristics of published and unpublished NCC cases

	<b>Published cases (n=211)</b>	<b>Unpublished cases (n=82)</b>	<b>Total (n=293)</b>

		n (%) <sup>1</sup>	n (%) <sup>1</sup>	n (%) <sup>1</sup>
Sex	Female	102/191 (53)	47/78 (60)	149/269 (55)
	Male	89/191 (47)	31/78 (40)	120/269 (45)
Age at diagnosis	Median age in years [IQR]	32 [21–47]	35 [26–46]	33 [23–47]
	Children (<18 years)	38/198 (19)	4/79 (5)	42/277 (15)
	Adults	160/198 (81)	75/79 (95)	235/277 (85)
Autochthonous cases		26/174 (15)	32/70 (46)	58/244 (24)
Travel/Migration <sup>2</sup>		148/174 (85)	38/70 (54)	186/244 (76)
Africa		43/148 (29)	16/38 (42)	59/186 (32)
Asia		33/148 (22)	22/38 (58)	55/186 (30)
Caribbean		11/148 (7)	1/38 (3)	12/186 (6)
Latin America		61/148 (41)	9/38 (24)	70/186 (38)
Middle East		3/148 (2)	0/38 (0)	3/186 (2)
Signs/ Symptoms	Epileptic seizures	109/199 (55)	49/68 (72)	158/266 (59)
	Focal onset seizures	20/73 (27)	2/33 (6)	22/106 (21)
	Generalised onset seizures	53/73 (73)	31/33 (94)	84/106 (79)
	Headache	92/187 (49)	40/68 (59)	132/255 (52)
	Other neurological signs/symptoms	90/198 (46)	34/35 (97)	125/233 (54)
Serology (Serum/CSF)	Antigen or antibody positive	89/122 (73)	42/62 (68)	131/184 (71)
	Antibody positive	63/75 (84)	NA	63/75 (84)
	Antigen positive	7/9 (78)	NA	7/9 (78)
Neuroimaging /EEG	CT	151/192 (79)	36/39 (92)	187/231 (81)
	CT with contrast	58/78 (74)	NA	58/78 (74)
	Only CT	28/151 (19)	2/36 (6)	30/187 (16)
	MRI	165/194 (85)	39/41 (95)	204/235 (87)
	MRI with contrast	127/132 (96)	NA	127/132 (96)
	Only MRI	40/165 (24)	3/32 (9)	43/197 (22)
	Soft tissue x-ray	39/177 (22)	NA	39/116 (22)
	EEG	24/167 (14)	NA	24/125 (14)
Results on neuroimaging	Single lesion	65/179 (36)	26/43 (60)	91/222 (41)
	Multiple lesions	114/179 (63)	17/43 (40)	131/222 (59)
	Viable cysts	177/198 (89)	57/61 (93)	234/259 (90)
	Single	75/167 (45)	28/41 (68)	103/208 (50)
	Multiple	92/167 (55)	13/41 (32)	105/208 (50)
	Enhancing cysts	99/145 (68)	9/13 (69)	108/158 (68)
	Cysts with scolex	80/134 (60)	5/9 (56)	85/143 (59)
	Calcifications	69/182 (38)	9/12 (75)	78/194 (40)
	Single	6/60 (10)	NA	6/60 (10)
	Multiple	54/60 (90)	NA	54/60 (90)

	Perilesional edema	83/156 (53)	41/62 (66)	124/218 (57)	
	Hydrocephalus	38/180 (21)	NA	38/180 (21)	
	Vesicular stage <sup>3</sup>	41/144 (29)	NA <sup>5</sup>	41/144 (29)	
	Colloidal/granular nodular stage <sup>3</sup>	82/144 (57)	NA <sup>5</sup>	82/144 (57)	
	Calcified stage <sup>3</sup>	21/144 (15) <sup>4</sup>	NA <sup>5</sup>	21/144 (15) <sup>4</sup>	
Cyst(s)/Calcification(s) location <sup>2</sup>	Cerebral	175/185 (95)	56/56 (100)	231/241 (96)	
	Intraparenchymal	149/175 (85)	52/56 (93)	201/231 (87)	
	Intraventricular	40/175 (23)	4/56 (7)	44/231 (19)	
	Subarachnoid	23/175 (13)	3/56 (5)	26/231 (11)	
	Spinal	16/168 (11)	2/3 (67)	18/171 (12)	
	Intra-medullary	3/16 (19)	1/1 (100)	4/17 (24)	
	Extra-medullary	13/16 (81)	0/1 (0)	13/17 (76)	
	Extra-neural	23/138 (17)	7/12 (58)	30/150 (20)	
	Treatment	Surgery	64/162 (40)	7/14 (50)	71/176 (40)
		Pathological examination	49/56 (88)	5/6 (83)	54/62 (87)
	Anthelmintic therapy	136/175 (78)	55/62 (89)	191/237 (81)	
	Praziquantel and Albendazole	19/136 (14)	9/55 (16)	28/191 (15)	
	Praziquantel only	14/136 (10)	3/55 (5)	17/191 (9)	
	Albendazole only	103/136 (76)	43/55 (78)	146/191 (76)	
	Corticosteroids	111/166 (67)	44/47 (94)	155/213 (73)	
	Prednisolone/Prednisone	30/78 (38)	9/17 (53)	39/95 (41)	
	Dexamethasone	48/78 (62)	17/17 (100)	65/95 (68)	
	Anti-epileptic treatment <sup>2</sup>	76/162 (47)	32/35 (91)	108/197 (55)	
	Carbamazepine/Oxcarbazepine	13/42 (31) <sup>2</sup>	NA	13/42 (31) <sup>2</sup>	
	Phenobarbitone	3/42 (7) <sup>2</sup>	NA	3/42 (7) <sup>2</sup>	
	Phenytoin	6/42 (14) <sup>2</sup>	NA	6/42 (14) <sup>2</sup>	
	Valproic acid	8/42 (19) <sup>2</sup>	NA	8/42 (19) <sup>2</sup>	
	Lamotrigine	3/42 (7) <sup>2</sup>	NA	3/42 (7) <sup>2</sup>	
	Levetiracetam	11/42 (26) <sup>2</sup>	NA	11/42 (26) <sup>2</sup>	
	Clobazam	1/42 (2) <sup>2</sup>	NA	1/42 (2) <sup>2</sup>	
Outcome	Cured	68/155 (44)	NA	68/155 (44)	
	Improved	71/155 (46)	NA	71/155 (46)	
	No change, deteriorated	11/155 (7)	NA	11/155 (7)	
	Death	5/155 (3)	NA	5/155 (3)	

360 1 Counted, if explicitly stated, visible on imaging or if it could be inferred; the denominator varies between variables.

361 2 More than one answer possible; hence the sum of the percentages can exceed 100%.

362 3 Stage of the cyst: Degenerative if at least one cyst is in the degenerative stage; vesicular and calcified stage if all cysts are in the respective stage. Hence, the numbers of viable  
363 cysts/calcifications do not match the results by stage.

364 4 This is the proportion of patients with only calcified stage among those with detailed information of the stage of the cysts. The overall proportion of patients with only calcified cysts is  
365 10%.

366 5 Due to limited information, no analyses were conducted for unpublished cases.

367 IQR Interquartile range  
368 NA Not available  
369

370 **TABLE 2.** Other neurological signs/symptoms

<b>Neurological sign/symptom</b>	<b>Cases with neurological signs/symptoms (N=125)<sup>2</sup></b>
Unsteadiness of gait <sup>1</sup>	33 (33%)
Cognitive impairment	26 (21%)
Impaired consciousness	25 (20%)
Impaired vision <sup>1</sup>	24 (19%)
Limb paresis	21 (17%)
Speech difficulties <sup>1</sup>	14 (11%)
Cranial nerves lesions	14 (11%)
Limb Ataxia	9 (7%)
Meningism <sup>3</sup>	9 (7%)
Impaired sensation <sup>1</sup>	9 (7%)
Vertigo	8 (6%)
Limb spasticity	6 (5%)
Bladder dysfunction	4 (3%)

371

372 <sup>1</sup> Speech difficulties, unsteadiness of gait, impaired vision and impaired sensation were not specified in more detail in the case descriptions, therefore origin (peripheral/central,  
373 cerebral/cerebellar etc.) remains unclear.

374 <sup>2</sup> Most symptomatic cases reported more than one neurological sign/symptom, therefore column totals are larger than N=125 or 100%, respectively.

375 <sup>3</sup> Signs/symptoms described as: "meningeal signs, neck stiffness, meningism, neck rigidity"

**TABLE 3.** Neurological signs/symptoms stratified by cyst(s) location, number, radiological characteristics and stage<sup>1</sup>

			Epileptic seizures		Headache		Other neurological signs/symptoms <sup>5</sup>	
			n (%)	p	n (%)	p	n (%)	p
Cyst(s) location	Intraparenchymal cyst(s)	Yes	129/196 (66)	p<0.001	96/183 (52)	p=0.31	95/166 (57)	P=0.16
		No	5/36 (14)		24/38 (63)		30/39 (77)	
	Intraventricular cyst(s)	Yes	10/40 (25)	p<0.001	32/44 (73)	p=0.008	29/41 (71)	p=0.14
		No	121/188 (64)		84/173 (49)		80/161 (50)	
	Subarachnoid cyst(s)	Yes	9/25 (36)	p=0.05	18/24 (75)	p=0.04	17/24 (71)	p=0.57
		No	119/201 (61)		96/191 (50)		91/177 (51)	
	Only intraparenchymal <sup>2</sup>		73/102 (72)		39/91 (43)		30/101 (30)	
Only intraventricular <sup>2</sup>		2/13 (15)		12/14 (86)		10/14 (71)		
Only subarachnoid <sup>2</sup>		1/7 (14)		4/7 (57)		5/7 (71)		
Cyst number	Single		64/89 (72)	p<0.001	35/81 (43)	p=0.14	34/71 (53)	p=0.60
	Multiple		62/126 (49)		68/124 (55)		63/119 (48)	
Concomitant findings	Ring enhancement <sup>3</sup>	Yes	67/106 (63)	p=0.06	41/94 (44)	p=0.08	44/100 (44)	p=0.78
		No	21/48 (44)		30/48 (62)		25/50 (50)	
	Perilesional edema <sup>3</sup>	Yes	87/121 (72)	p<0.001	50/113 (44)	P=0.009	48/99 (48)	p=0.60
		No	41/91 (45)		57/89 (64)		55/88 (62)	
	Hydrocephalus	Yes	7/36 (19)	p<0.001	26/39 (67)	p=0.04	34/40 (85)	p<0.001
		No	90/140 (64)		59/128 (46)		52/139 (34)	
Stage	Vesicular stage		11/32 (34)		22/34 (65)		16/34 (47)	
	Degenerative stage <sup>4</sup>		82/116 (71)		47/107 (44)		46/95 (48)	
	Calcified stage		15/23 (65)		9/24 (38)		10/24 (42)	

<sup>1</sup> Only patients with detailed information on stage of the cyst(s) were included (published and unpublished cases). Data refer to the known information on epileptic seizures, headache and other neurological signs/symptoms by group categorised in the left hand column (e.g. intraparenchymal cysts). "n" gives the number of patients with the respective sign/symptom. For example: of the 196 patients with intraparenchymal cysts, 129 (66%) presented with epileptic seizures. Apart from the category "stage", more than one criterion was possible per category.

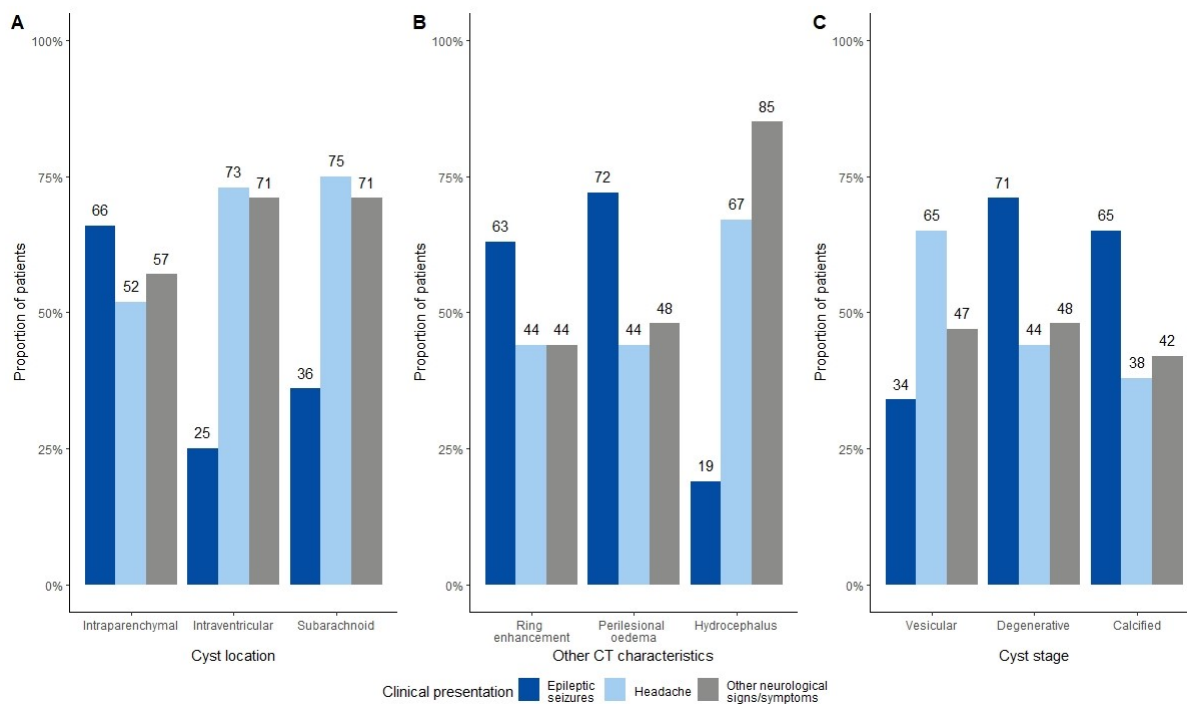
<sup>2</sup> Patients with cysts at various locations (intraparenchymal, intraventricular, subarachnoid) were excluded.

<sup>3</sup> Ring enhancement/perilesional edema around at least one lesion



4 Degenerative stage: colloidal or granular nodular stage

5 Detailed list of other neurological findings can be found in Table 2



347 **FIGURE 2.** Clinical presentation by cyst location (A), neuroimaging findings (B) and cyst stage (C). For cyst  
 348 location and CT characteristics more than one criterion was possible. \* Stage of the cyst: Degenerative if at least one  
 349 cyst is in the degenerative stage; vesicular and calcified stage if all cysts are in the respective stage.

351 **Discussion**

352 In this study, we present demographic and clinical details of almost 300 patients with NCC in the  
 353 European context, based on a systematic literature search, including grey literature, in addition to  
 354 unpublished NCC cases collected from colleagues through the European network CYSTINET. We  
 355 report several novel observations as well as observations that are in line with previously published  
 356 literature [16–19,33,34].

358 *Origins of infection*

360 We were able to show that most patients treated for NCC in Europe are migrants from countries  
 361 endemic for *T. solium*. However, confirming the exact region of origin of infection is a challenge, as NCC  
 362 often becomes symptomatic only several years after first exposure. There seems to be only few regions  
 363 in Europe where the full lifecycle of *T. solium* is still present and in those regions, autochthonous NCC  
 364 cases can still occur [18]. Our search yielded relatively more autochthonous cases among unpublished  
 365 NCC cases compared with published cases. This is likely due to reporting differences between countries  
 366 of published and unpublished cases, e.g. especially the reported cases in Romania were mostly  
 367 unpublished. The number of autochthonous cases reported should be interpreted with caution, as

368 routine travel history reported usually included only a few non-standardized questions and did not  
369 include a detailed epidemiologic interview. Also, it only requires one tapeworm carrier to infect other  
370 people with cysticercosis – for autochthonous cases, this person could either be another local person but  
371 could also be a migrant which makes it difficult to trace back the source of infection. In our review, we  
372 also found a considerably lower proportion of autochthonous cases compared with a review on  
373 cysticercosis in Europe, by Zammarchi et al. in 2013, which found 62% of cysticercosis cases being  
374 autochthonous cases which may suggest improvements in the disruption of the *T. solium* lifecycle [16].

375

#### 376 *Clinical characteristics and related findings on neuroimaging*

377

378 The sites and stages of cysts influences how patients with NCC present. With regards to site, most  
379 symptomatic patients with intraparenchymal cysts presented with epileptic seizures compared to  
380 patients with extraparenchymal cysts who were more likely to present with non-specific symptoms like  
381 headache. Patients with extraparenchymal cysts were significantly older at diagnosis. This may be  
382 because patients infected at an older age were more likely to develop extraparenchymal NCC e.g. due  
383 to comorbidities and immunological factors or because symptoms were less specific and therefore  
384 diagnosis was made at a later stage [35]. These findings, i.e. patients with extraparenchymal lesions  
385 being older and showing more non-specific signs/symptoms, concur with NCC data published from  
386 Mexico [5]. The number of patients with autochthonous cases was rather low so it was difficult to draw  
387 conclusions on disease presentation in comparison to patients you were infected in Latin America, Asia  
388 or Africa.

389 Ring enhancing lesions and those with perilesional edema also seemed to predispose to presentation  
390 with epileptic seizures. However, although epileptic seizures occurred frequently under those  
391 conditions, around half of the cases were also accompanied by other neurological signs and symptoms,  
392 including headache. The latter, however, was frequently reported in extraparenchymal NCC  
393 (intraventricular and subarachnoid NCC) and when hydrocephalus was present. Looking at cyst stage,  
394 headache prevailed in the vesicular stage, where little to no inflammation was evident on radiology, the  
395 pathophysiology of headache under these conditions is not clear (ASW unpublished data), whereas  
396 epileptic seizures were the predominant neurological symptom during the degenerative stage of the  
397 cyst(s)

398 It is well established that clinical manifestations of NCC can vary from completely asymptomatic  
399 infection (54% of NCC patients in a study of Monteiro de Almeida et al. [36]) to severe disease and death  
400 [2,37]. The major determinants of the characteristics of symptomatic NCC are the number of cysts, their  
401 location, their stage and the degree of inflammation [35,38]. It has been shown previously that NCC can  
402 mimic almost any neurological disorder [39], but to date neurological signs/symptoms other than

403 epileptic seizures or headache are thought to occur in the minority of patients [1]. In concordance with  
404 previous studies, we found a high proportion of epileptic seizures among NCC patients [1,16,40].  
405 However, in our study of European presentations, we found that more than half of all NCC patients  
406 presented with other neurological signs/symptoms. This may reflect some publication bias, as unusual  
407 case presentations more likely come to the attention of a clinician and be published than patients  
408 presenting with well-known signs/symptoms of NCC. Also, patients presenting in low-income and  
409 middle-income countries may be more likely to present later to health services than in high-income  
410 countries, often with more severe symptoms and subtle neurological signs and symptoms may not be  
411 identified or recorded as consistently.

412 Regarding publications included in our study, the classification of neurological signs/symptoms other  
413 than epileptic seizures and headache was challenging as description of signs/symptoms could be vague  
414 and their origin remaining uncertain (see Table 3). Remarkably, in our study, impaired cognitive  
415 function appeared to be one of the most frequent other neurological signs. In the included publications,  
416 EEG was rarely performed and predominantly only when epileptic seizures were present. Therefore,  
417 non-convulsive seizures that could account for impaired cognitive function may not have been  
418 identified. A high proportion of cognitive decline (87.5%), dementia (12.5%) or altered mental state  
419 (28%) in NCC patients has also been observed in previous studies [1,41,42].

420

#### 421 *Special considerations*

422

423 According to previously published studies, between 1.5% and 3% of NCC patients are estimated to have  
424 spinal cysts [43–48]. The true proportion is likely higher than 3% as spinal imaging is not routinely  
425 performed. In our dataset, this proportion was higher (18/171; 10.5%). For example, a study in Peru  
426 found spinal cysts in 17 of 28 (61%) patients with basal subarachnoid cysts, supporting a  
427 recommendation to perform spinal imaging in all patients with basal subarachnoid NCC [49]. In our  
428 dataset, 5/27 (19%) patients with subarachnoid NCC also had spinal cysts, while not all 27 patients had  
429 cysts in the basal subarachnoid space. Combined cerebral and spinal imaging was not often done, or at  
430 least not often reported in our European cases and may result in an underestimation of the true burden  
431 of spinal NCC in the context of subarachnoid NCC in our study population.

432 In addition to spinal NCC, we would like to highlight the identification of perilesional edema around a  
433 calcification. In our study population, we found one patient where perilesional edema around a  
434 calcification was present. Until recently, calcifications were considered inactive in terms of immune  
435 response. However, it seems that calcified cysticerci can temporarily release residual antigen that may  
436 trigger an immune response [50–52]. Why this happens is still not entirely clear. It could make treatment  
437 in patients with concomitant viable cysts and calcifications challenging, as epileptic seizures may still

438 occur after cyst resolution (after anthelmintic therapy or spontaneously) and therefore discontinuation  
439 of AED after resolution of viable cysts in patients with concomitant intermittent perilesional edema  
440 around calcifications may carry risks. Clinicians should be aware that this may occur. Perilesional  
441 edema around calcifications should not be treated with anthelmintic drugs and corticosteroid therapy  
442 is not routinely recommended [53].

443

#### 444 *The value of serology and neuroimaging*

445

446 In our study, serological testing of any kind for *T. solium* in CSF and/or serum was reported in only 61%  
447 of cases. It is likely that serological testing was performed as confirmation after neuroimaging, however,  
448 from our dataset we could not establish why serological testing was performed in some cases and not  
449 others, nor the time point when it was performed in relation to presentation. Serological detection of *T.*  
450 *solium* antigen and antibodies is not yet widely available throughout Europe. There are only a few  
451 commercial tests available on the market, and these are often supplied in a form that is not always  
452 suitable for laboratories that see only occasional cases. In addition, in-house assays are not easy to  
453 implement and validate for laboratories due to a lack of well-defined reference sera. A commercial  
454 antigen test has only been available in the last few years. Until then, in-house assays relied on  
455 monoclonal antibodies which were not readily available. The low number of antigen tests reported is  
456 particularly striking, as there is good evidence that antigen follow-up is useful in the context of  
457 therapeutic outcome monitoring in symptomatic active NCC cases. Here, clear and easily available  
458 guidelines for *T. solium* serological testing seem desirable. Although antigen testing was performed  
459 much less frequently than antibody testing, it showed higher sensitivity. However, it must be taken into  
460 account that the proportion of patients with active stage lesions was rather high which may have  
461 influenced the sensitivity of the antigen ELISA. Still, one-third of the confirmed NCC cases, more so  
462 those with single lesions, were not positive in any serological assay which demonstrates the need for  
463 establishing a standardized approach to immunodiagnostic testing in European laboratories.

464 Regarding neuroimaging, a combination of MRI and CT imaging is recommended in the Clinical  
465 Practice Guidelines for the Diagnosis and Treatment of NCC by the Infectious Diseases Society of  
466 America (IDSA) and the American Society of Tropical Medicine and Hygiene (ASTMH) guidelines and  
467 in the WHO guidelines on management of *Taenia solium* neurocysticercosis, because MRI has a higher  
468 sensitivity for detecting active NCC lesions and CT imaging has a higher sensitivity for detecting  
469 calcifications [53,54]. In our study, a third of all patients only had one type of neuroimaging performed,  
470 which may have lead to an underreporting of both active and inactive NCC lesions.

471

#### 472 *Treatment of symptomatic patients with neurocysticercosis*

473

474 In the IDSA/ASTMH guidelines, different recommendations for anthelmintic treatment are given for  
475 different manifestations of NCC [53]. Surgical removal and/or ventricular shunting in the case of  
476 increased intracranial pressure is recommended for patients with intraventricular cysts. For patients  
477 with NCC-related hydrocephalus or diffuse cerebral edema, corticosteroid therapy without  
478 anthelmintics is recommended. For patients with one or two intraparenchymal cysts, 15mg/kg/day  
479 albendazole is advised for 10-14 days with treatment extended if no effect in terms of cyst resolution  
480 is shown; for patients with more than two viable intraparenchymal cysts it is recommended to combine  
481 albendazole with praziquantel 50mg/kg/day is recommended [53]. Generally, treatment  
482 recommendations do not differ for children and adults, apart from an adjustment of anthelmintic drug  
483 dosage. In our dataset, which includes many cases that were diagnosed and treated before the above  
484 guidelines were published in 2018, approaches to treatment were extremely varied and longer durations  
485 and higher dosages were given to patients with more severe manifestations or to non-responders. This  
486 reflects the importance of having clear consensus guidance for treatment in order to improve safety and  
487 efficacy, and to better understand treatment outcomes.

488 In our European population, corticosteroids were mostly administered for the same duration as  
489 anthelmintic therapy although surprisingly, many patients received corticosteroids for shorter time  
490 than anthelmintics. This had been recommended previously [7,8], but is nowadays not recommended.  
491 However, this could also be due to the fact that concurrent steroid therapy was not consistently  
492 reported. For corticosteroid therapy, the IDSA/ASTMH guidelines recommend steroids for the entire  
493 duration of the anthelmintic therapy and even starting 3 to 4 days in advance. Generally,  
494 dexamethasone and prednisolone are recommended. For dexamethasone, a dosage of 0.1mg/kg/d is  
495 used for patients with intraparenchymal cysts and up to 0.2mg/kg/day for patients with  
496 extraparenchymal cysts in the basal cisterns or Sylvian fissure. Successful treatment is contingent on  
497 finding the right steroid dose which on the one hand prevents side effects of anthelmintic therapy (i.e.  
498 epileptic seizures, severe headache or increased intracranial pressure), but on the other hand also does  
499 not suppress the effect of anthelmintic drugs. This is particularly important when treating with a  
500 combination therapy of albendazole and praziquantel, as there are enzymatic interactions with  
501 dexamethasone, which may lower plasma levels of praziquantel [55]. The IDSA/ASTMH guidelines do  
502 not specify whether steroids should be given even after anthelmintic therapy has ended, but do  
503 recommend that steroids should be tapered if they have been given for more than two weeks [53]. It is  
504 important to note that the cysticidal effects of anthelmintic therapy may last beyond the end of the  
505 treatment cycle, especially with combination therapy with albendazole and praziquantel, and reducing  
506 steroid doses over a longer period may be advisable in these cases, at least until disappearance of any  
507 perilesional oedema.

508 We found different AEDs used in patients treated in our European population. Whilst AEDs are  
509 recommended for all patients with epileptic seizures according to IDSA/ASTMH guidelines, no  
510 recommendation on the type is given; choice should be based on local availability, cost, drug-drug  
511 interactions, and potential side effects [53].

512

### 513 *Recommendations for diagnosis and treatment of neurocysticercosis*

514

515 With this publication we would like to raise the awareness of NCC among clinicians working in  
516 European countries. While a travel history to or from endemic countries may support a suspected  
517 diagnosis of NCC, the absence of a relevant travel history does not exclude the presence of NCC. In  
518 addition, NCC remains predominantly a neuroradiological diagnosis, which means that performing a  
519 neuroradiological examination, preferably CT and MRI combined, should be the first priority. At  
520 present, serological testing plays only a supportive role in the case of suspicious cerebral lesions on  
521 imaging. In addition, the possibility of spinal NCC and temporary perilesional edema around  
522 calcifications must be considered and neuroimaging tailored to answer these questions.

523 With a few exceptions, mainly regarding the availability of serological tests and some drugs, until  
524 guidelines tailored to the European context may become available, the IDSA/ASTMH guidelines can,  
525 for the time being, be applied to the European context, although there may be only limited access to  
526 MRI in some areas of Europe. Based on our experience with the IDSA/ASTMH guidelines, we  
527 recommend that the next update reconsiders the treatment guidelines for (co-existing) hydrocephalus  
528 and the length and dose of concomitant steroid therapy in combination therapy with albendazole and  
529 praziquantel.

530

### 531 *Neurocysticercosis in the context of migration and travel* 532 *medicine*

533 We were able to demonstrate that neurocysticercosis occurs in Europe albeit most of the cases occur  
534 among migrants and travellers. With increasing travel and migration, an increase in NCC cases in  
535 Europe can also be expected although globally, the number of NCC cases are decreasing [56]. This  
536 phenomenon has been described for other neglected tropical diseases before [57]. Migrants often face  
537 inequalities in access to healthcare which hampers diagnosis of infectious diseases, and which may have  
538 been even stronger during the COVID-19 pandemic [58,59]. There has been increasing effort by the  
539 International Society on Travel Medicine to promote migrant health and to bring to attention diseases  
that more commonly occur among migrants. Hence, this study can also serve this purpose.

540

### 541 *Strengths and limitations*

542

543 The strength of the current paper lies in the combination of original patient data from unpublished NCC  
544 cases combined with patient characteristics gathered through an exhaustive systematic literature search  
545 including grey literature, in several major European languages. Detailed clinical and demographic  
546 characteristics of NCC patients diagnosed and treated in Europe have not previously been so  
547 comprehensively summarized in one document, and this study will therefore be of value to any clinician  
548 in Europe engaged in the diagnosis or treatment of NCC. Europe is a unique setting for NCC as *T. solium*  
549 is not highly prevalent while diagnostic facilities, particularly neuroimaging, are widely available.  
550 Regardless, a number of important limitations should be acknowledged. Full text access to the huge  
551 number of identified NCC related publications worldwide was not possible, and neither was it possible  
552 to acquire access to all country-specific publications. Inherent publication biases are, however, likely to  
553 be of greater importance: a) only selected cases being published, and b) often no detailed clinical  
554 information being provided in case reports. Also, sharing of unpublished cases, as well as grey  
555 literature, depends on research interests and cooperation, and identifying all unpublished NCC cases in  
556 Europe exceeded our research capacity. Thus, the data presented in this study are prone to bias,  
557 potentially towards more over-representation of more unusual cases. In addition, assessment of  
558 outcomes is limited by lack of standardisation of timing at which the outcome may have been  
559 determined.

560

## 561 **Conclusions**

562 The synthesis of knowledge of and information about NCC in the European context contained in this  
563 publication is unprecedented. NCC represents a rare disease for European clinicians and hence clinical  
564 familiarity outside specialist centres is most likely scarce. The current publication contributes to a better  
565 understanding of the origin of infection with *T. solium*, clinical characteristics, diagnosis and treatment  
566 of patients suffering from NCC in Europe. These data highlight that NCC causes considerable morbidity  
567 in patients diagnosed in Europe and that overall treatment outcomes can be improved. In some  
568 patients, NCC is a cause of death. Due to the complexity of the life cycle of *T. solium* and the latency  
569 until signs and/or symptoms appear, reliable multi-/interdisciplinary data are needed to establish  
570 standardized context-specific evidence-based management guidelines for practising clinicians in  
571 Europe and beyond.

572

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583

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